# Acid sanitizing and cleaning compositions containing protonated carboxylic acids

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A sanitizing composition comprising at least one aliphatic short chain antimicrobially effective C5 to C14 fatty acid or mixture thereof, at least one carboxylic weak acid and a strong mineral acid which may be nitric or a mixture of nitric and phosphoric acids.

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# Acid sanitizing and cleaning compositions containing protonated carboxylic acids

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#### FIELD OF THE INVENTION

[0002] The present invention relates to acid sanitizing and/or cleaning compositions comprising arithmicrobially effective S3 to C antiboxylic acids. The present invention is directed to both concentrates and to water diluted use solutions.

BACKGROUND OF THE INVENTION [0003] Periodic cleaning and sanitizing in food, drink, pharmaceutical, cosmetic and similar processing industries; in food preparation and service businesses; in health and day care facilities; and, in hospitality establishments are a necessary practice for product quality and public health. Residuals left on equipment surfaces or contaminants found in the process or service environment can harbor and nourish growth of subsequent processed product or critical contact surfaces. Protecting the consumer against potential health hazards associated with pathogens or toxins and maintaining the quality of the product or service requires routine cleaning of residuals from surfaces and effective sanitation to reduce microbial populations. [0004] Visual inspection of the equipment cannot ensure that surfaces are clean or free of microorganisms. Antimicrobial treatments as well as cleaning treatments are therefore required for all critical surfaces in order to reduce microbial population to safe levels established by public health regulations. This process is generally referred to as sanitizing. The practice is of sanitation is particularly of concern in food process facilities wherein the cleaning treatment is followed by an antimicrobial treatment applied upon all critical surfaces and environmental surfaces to reduce the microbial population to safe levels established by ordinance. A sanitized surface is, as defined by the Environmental Protection Agency (EPA), a consequence of a process or program containing both an initial cleaning and a subsequent sanitizing treatment which must be separated by a potable water rinse. A sanitizing treatment applied to a cleaned food contact surface must result in a reduction in population of at least 99.999% (5 log) for specified microorganisms as defined by the "Germicidal and Detergent Sanitizing Action of Disinfectants", Official Methods of Analysis of the Association of Official Analytical Chemists, paragraph 960.09 and applicable sections, 15thEdition, 1990 (EPA Guideline 91-1). [0005] The antimicrobial efficacy of sanitizing treatments is significantly reduced if the surface is not

locoty in the attentional emiscry to statuting understasts significantly records in the statistics and seasons absolutely free of soil and other containments prior to the santitizing step. The presence of reducial flood and/or mineral deposits inhibit santitizing treatments by acting as physical barriers which shield microorganisms high within the organic or inorganic layer from the microbidice. Furthermore, chemical interactions between the microbidide and certain contaminants can disrupt the killing mechanism of the microbidide.

[0005] With the advent of automated clean-in-place and samitize-in-place systems, the need for disassembly has been diminished, and cleaning and samitizing have become much more effective. However, modern food inclustres still rely on a smitzers to compensate for design deficiencies or operational limitations in their cleaning programs and the probability of very small residual amounts of organic and integrate so tests and higher performance expectations, scattlers treatments must also comply with the increasing demand for safer, less corrective, more environmentally friendly compositions.

[0007] According to the U.S. Center for Disease Control and Prevention, tood poleoning in calendar year 2000 resulation in 5000 dastites, 250,000 hospitalizations and 750,000,000 illnesses. The need voists for improved santizing freshments to destroy pathogens and food spoilage microorganisms resistant to conventional reshments within the tool gathering, food processing, and food serving inclusivies. A further complication is that the list of approved microbiotidal agents has continued to decrease due to acute and chronic human toxicity of some microbiotidal agents, and to their environmental persistence in weter

[0005] Artimicrobially active adds have been used in santitizing operations. For instance, U.S. Pat. No 40,400 describes a santilizing operation comprising solishest, short chain first adds, a hydrotrope or solubilize for the fatty adds, and a hydrotrope-compatible add, and U.S. Pat. No. 5,330,760 describes fally add santilizer concentrates and dilute final solutions within indude individual amounts of germaticeally effective fatty add, hydrotrope, a strong add group consisting of phosphoric add and sulfuric add or mixtures thereof sufficient to lover the pit of the final solutions abd out 1-5, and a concentrate stabilizing weak add component selected from the group consisting of propionic, butyric and valeric adds and mixtures thereof a final solutions.

[0009] Protonated carboxylic acids offer broad spectrum antimicrobial activity against gram-positive and gram negative bacteria, persistent blocklal activity in the presence of organic and inorganic soils and residual blocklal and inhibitory activity. They combine both acid for mineral deposit control and santitizer for antimicrobial effect his one treatment solution.

[0010] However, one problem associated with the use of protonated carboxyle acid sanifizers is poor use distribution phase satisfity, particularly at lower water temperatures of 40(tigo.) 5(diceg.) F. 50(tigo.) F. 6118 more carboxyle acids having alkyl chains containing 5 or more action atoms, are typically characterized as water insociation and can oil out or precipitate from solution as a gelatinous localization. Solve the process of the sanification is intended to sanificat, such as opportunity and sanification of the sanification of the

#### efficacy

[0012] Organic hydrotropies or coupling agents, such as low molecular weight suffonates, may be employed to increase the solubility and miscbility of the longer chain fathy adds with water and inorganic salts both in concentrated and in diluted use solutions. Again, the solubility appears to diminish at sustained lower water temperatures with the result being phase separation.

[0013] There remains a need in the art for an improved blocidal composition which utilizes a carboxylic fatty acid which has high antimicrobial efficacy, has good phase stability, exhibits low toxicity, and is not detrimental to the environment.

SUMMARY OF THE INVENTION

[0014] Surprisingly, the compositions of the present invention exhibit excellent phase stability both in concentrated form and in diluted use solutions, and in particular, they exhibit excellent phase stability in tow temperature water diluted use solutions. Even more surprisingly, the stability is improved in the presence of nitric acid.

month The sanitizing and/or dearing compositions of the present invention, in both concentrated and inbulled use solution, include an antimicrobally effective bort chain tells acid, a shorter chain weak carboxylic acid, and a strong mineral acid. The ebroter chain weak carboxylic acid functions as a solvent for the antimicrobally effective as a solvent for the antimicrobally effective acid functions as a solvent for the antimicrobally effective short chain fally acid. In concentrated form, the compositions also desirably contain an organic hydrotrope. [017] In some membediments, the entimicrobally effective short chain fally acid is a CS to C4 fattly acid,

[0017] in some embodiments, the antimicroscality effects short data may add its a C.5 to C.14 starty acid, and more suitably CG to C.10 starty acid, or some mixture thereof, the shorter chain weak carboxyfex acid is a only more chain to the control of the contr

[0.119] Additionally, the composition may optionally include at least one arionic and/or nonionic surfactant, in some embodiments, a nonionic surfactant is sutistly employed to improve surface wetting, soil removal, and so forth. It may also function to improve the solubility of the felty acids at use dilutions. [0.002] The arithmicrosality infective effective short chain fatty acid is useful from about 3 w-% to about 12

int-8 of the concentrate, and more suitably from about 5 wi-9k to about 10 wi-9k of the concentrate, in one perticular embodiment, the concentrate includes a blend of two fatty acids.
(IOC21) The weak carboxylic add is useful from about 5 wi-9k to about 50 wi-9k of the concentrate, and more suitably from about 10 wi-9k to about 40 wi-9k of the concentrate, in one particular embodiment, the weak carboxylic acid acts as a solvent for the carboxylic acid acts as a solvent for the

antimicrobially active short chain fatty acid.

(0022) The strong mineral acid is useful from about 5 wt-% to about 50 wt-% of the concentrate, and more suitably about 15 wt-% to about 40 wt-% of the concentrate. In some embodiments, the strong mineral acid is nitric which is useful from about 5 wt-% to about 50 wt-% of the concentrate, and more suita concentrate. If phosphoric acid is employed, it is useful from 5 wt-% a bout 40 wt-% of the concentrate, and more

suitably about 10 wt-% to about 35 wt-% of the concentrate.

[0023] Suryrisingly, the arthricoblaily active short chain fathy acti is stable in nitric acid.
[0024] The compositions may further comprise optional ingredients including urea for stabilization of nitric acid, and a surfactant component. The surfactant component may include one or more surfactants. In some embodiments, an anionic or nonionic surfactant may be optionally added at level of 0.1 wt. % to about 50 wt.% of the concentrate, even more suitably about 0.25 wt.% to about 40 wt.% of the concentrate, even more suitably about 0.4 wt.% of the concentrate, even more suitably about 1.4 wt.% of well on the order of the order order of the order order

alkystulforate. [0029] The compositions may be disuted with water at any ratio whatsoever, but typically the ratio is between about 1:100 parts of the concentrate to water to about 1:1500 parts of the concentrate to water. This is referred to as a use distinct, or every typical use distinct is about 1 conce of concentrate to about 6 gallons of water which is a ratio of about 1:768 parts of the concentrate to water. [0027] The compositions of the present invention find utility as both satisficiary and disinfecting compositions.

[0.027] The compositions of the present invention find utility as both santitizing and disinfecting compositions as well as chemical compositions, and are useful for both hard and soft surface sentitizing and disinfecting in farm operations, food processing operations, institutional food preparation and serving areas, health care and child care facilities as well as any other number of contact sensitive environments. The compositions exhibit high antimicrobial efficacy while having low toxicity, are not detrimental to the environment, and do not contaminate food stuffs.

[0028] The compositions also find utility for use as one-step cleaning/sanitizing compositions and disinfectants in which the composition cleans and sanitizes simultaneously.

### BRIEF DESCRIPTION OF THE FIGURES

[0029] FIG. 1 is a bar graph illustrating results of a foaming evaluation conducted for example 24 which was compared to three commercially available sanitizing compositions.

# DETAILED DESCRIPTIONS OF THE PREFERRED EMBODIMENTS

[0030] While this invention may be embodied in many different forms, there are described in detail herein specific embodiments of the invention. This description is an exemplification of the principles of the invention and is not intended to limit the invention to the particular embodiments illustrated. [0031] The antimicrobial agents useful herein include those referred to generally in the art as acid-anionics including carboxylic acids having biocidal activity when in protonated form. These antimicrobial agents are typically classified as having low toxicity and as being environmentally friendly.

[0032] The term "short chain fatty acids" as used herein refer to those acids generally having about 5 to 14 carbon atoms, suitably about 6 to 12 carbon atoms, more suitably from about 6 to 10 carbon atoms and most suitably about 7-10 carbon atoms. In some embodiments of the present invention, a blend of a C9, fatty acid and a C10 fatty acid or a blend of nonanoic and decanoic acids is employed. [0033] The short chain fatty acids are useful from about 3 wt-% to about 12 wt-% of the concentrate and suitably about 5 wt-% to about 10 wt-% of the concentrate. In some embodiments of the present invention which employ a blend, nonanoic acid is utilized from about 2 wt-% to about 10 wt-% of the concentrate, suitably from about 3 wt-% to about 9 wt-%, and more suitably from about 4 wt-% to about 8 wt-% of the concentrate while decancic acid is utilized from about 0.5 wt-% to about 5 wt-%, suitably from about 0.5 wt-% to about 4 wt-% and more suitably about 1 wt-% 3 wt-% of the concentrate. The present inventors have found that when employing such a blend of nonanoic and decanoic acids, the phase stability appears to be improved when employing more nonanoic acid, and less decanoic acid. It is surmised that the shorter chain of the nonanoic acid provides increased solubility in water over the decanoic acid, while the decanoic acid provides increased antimicrobial efficacy over the nonanoic acid. Blending the two has been found to be particularly advantageous.

To with its about 40 with of the concentrate (COSQ) The storing add component of the form conditions is utilized to lower the pit in the final solutions to a midure of risks and component of the international CSA. The storing solid is suitably efform entire acid, or a midure of risks and phosphoric acids. Nitric acid is useful from about 6 with 50 about 50 with 50 about 50 with 60 the concentrate, and autility from about 15 with 50 about 40 with 60 about 50 with 50 of the concentrate. The first particular acids in useful from about 0 with 10 about 40 with 60 the concentrate and more suitably about 5 with 50 about 55 with 50 the concentrate. The first particular place acids of the presence invention have been found to be particularly state in the presence in the compositions of the presence invention have been found to be particularly state in the presence in the compositions of the presence invention have been found to be particularly state in the presence in the compositions of the present invention have been found to be particularly state in the presence of the passive surface layor. Statisness steel by marriarcharance of the passive surface layor. Statisness steel by marriarcharance of the passive surface layor. Statisness steel is corrossion resistant due to an oxide firm layer on the surface resulting from treatment with storing oxidizing agents such

[0398] Another problem often associated with the use of acid sentitizes is corrosion of stainless steel surfaces associated with high mineal waters or softened waters containing choldres which promote and accelerate corrosion of such surfaces. Strong codifizing acids with coldizing potential sufficient to passivate stainless are capable of reduction or eliminating such a problem.

(0037) A small amount of urea may be optionally employed in the compositions of the present invention. Organic degradation can occur in the presence of nitric acid by oxidation and nitritation mechanisms due to the presence and codiciting owner of nitrogen decides (NCQ) and integree tetrode (NCQ)-collectively enterred to as nitrogen persordor. Uncer may be added to react with the nitrogen persordor to reduce the returned to as nitrogen persordor. Uncer may be added to react with the nitrogen between the control of the nitrogen persordor. Uncertainty of the nitrogen has not personal personal personal nitrogen, but is autitably used from about 0.05 w/s to about 5 w/s, and more suitably at a level of about 0.1 w/s to about 1.0 w/s of the concentrate.

[0038] Surfaciants may also be optionally added to the compositions of the present invention for a variety of reasons including improved surface vetting by lowering the surface tension, improved soil or blofflim penetration, improved soil or blofflim penetration, removal and suspension of organic soils, enhancement of blockdal effect, characterization of loom profile, it. b. by the addition of low feating and high forestance coupling agent for the fatty add artiminizobial to mention a few. One skilled in the art will understand that some surfaciants or midurate of surfaciants seven one or more of these purposes better than others. The surfaciant or midurate of surfaciants seven one or more of these purposes better than others. The surfaciant or midurate of surfaciants seven one or more of these purposes better than others. The surfaciant or midurate of surfaciants seven one or more of these purposes better than others. The surfaciant or midurate of surfaciants seven one or more of these purposes better than others. The surfaciants or midurate of surfaciants seven the one intensity of the composition of the resolution of the surfaciant seven the rem include nonlonic, another onlone and cation is cartificants. Most suitably, the (0039) The surfaciants useful herein unique nonlonic and cation is cartificants.

[0039] The surfactants useful herein include nonionic, anionic and cationic surfactants. Most suitably, the surfactants employed include water soluble or water dispersible anionic or nonionic surfactants, or some combination thereof.

[0040] Useful anionic surfactants include, but are not limited to, those compounds having an hydrophobic group of C6-22 such as alky, alkylaryl, alteryl, acyl, long chain hydroxyalkyl, alkoxylated derivatives tread and so forth, and at least one water-solubilizing group of acid or salt from derived from sulmicin acid, sulfuric acid seter, phosphoric acid ester and carboxylic acid. The salt may be selected based on the specific formulation to which it is being addition.

[0041] More suitably, the anionic surfactants useful herein include, but are not limited to, sulfonated anionics such as alkyl sulfonates or disulfonates, alkyl anyl sulfonates, alkyl naphthalene sulfonates, alkyl diphenyl oxide disulfonates, and so forth.

[OO22] More particularly, the aninonic surfactants more suitable for use herein include, but are not limited by, those aninonic surfactants with: he arise or branched CFC-14 allybboranes suiforates, skilly an apphtitution suiforates, long chain all-knee suiforates, long chain all-knee suiforates, long chain particularly suiforates, suiforates including corresponding classificantes including 1-classe suiforates suiforates, skilly providents, skill

[0.043] Specific examples of anionic surfactants suitable for use herein include ally sulfonates such as 1cations sulfonate commercially available from a variety of including Stepan Co. in Northfield, II. under the tradersme of BIO-TERGE(PR) PAS-8; PILOT(R) L-45, a C11.5 allyberozene sulfonate (referred to as T4.57) from Pilot Chemical Co.; BIOSOF(PR) 500 and s130, non-neutralized internal sulforenzers estionic acids (referred to as "HLAS"), and S40, also an LAS, all from Stepan Company, DOWFAX(PR) anionic allylated (referred to as "HLAS"), and S40, also an LAS, all from Stepan Company, DOWFAX(PR) anionic allylated (referred to as "HLAS"), and S40, also an LAS, all from Stepan Company, DOWFAX(PR) anionic allylated (referred to as "HLAS"), and S40, also an LAS, all from Stepan Company, DOWFAX(PR) anionic allylated (referred to as "HLAS"), and S40, also an LAS, all from Stepan Company, DOWFAX(PR) anionic allylated (referred to as "HLAS"), and S40, also an LAS, all from Stepan Company, DOWFAX(PR) anionic allylated (referred to as "HLAS"), and S40, also an LAS, all from Stepan Company, DOWFAX(PR) anionic allylated (referred to as "HLAS"), and S40, also and LAS, all from S40, and set the set of the set of

(0.044) Examples of nonionic surfactants useful in the compositions of the present invention include, but are not limited to, the following classes:

[GO45] 1) polyoxyproxylene-polyoxylethylene block polymers including those made from propoxylation and/or ethoxylation of an initiator hydrogen compound such as propylene glyoc, ethylene glyoc, glyoran, trimethyldpropane, ethylenediamine, and so forth such as those sold under the tradename of PLURONIC (R) AND TETRONICRI available from BAST Corp.

[00:46] 2) condensation products of one mole of C8 to C18 branched or straight chain alkyl or dialkyl phenol with about 3 to about 50 moles of ethylene oxide such as those sold under the tradename of IGEPAL(R) available from Rhone-Poulenc and TRITOR(R) available from Union Carbide.

[0.047] 3) condensation products of one mole of a saturated or unsaturated, branched or straight C6 to C24 action-bit with about 3 to about 50 moles of ethylene oxide such as those odd under the tradename of NECHOL(R) available from Shell Chemical Cc. and ALFONIC(R) available from Condea Vista Cc.; [0.048] of condensation products of one note of saturated or unsaturated, burnarded or straight-chain C8 to 10.048] or condensation products of one note of saturated or unsaturated, burnarded or straight-chain C8 to tradename of NOPALCOL(R) from Herket Corp. and LIPO/PEG(R) from Lipo Chemicala, Inc.; and other alkanoic extens formed by condensation of catrolytic acids with glycordica, gyloreni, and polytyridir control of the contr

#### alcohols;

[0049] 5) surfactants produced by the sequential addition of ethylene oxide and propylene oxide to ethylene splot, ethylenediamien which result in a hydrophie with hydrophic blobbed (i.e. propylene oxide) at the terminal ends (the hydrophic) and hydrophobb (blobs are reversed) of each molecula weighing from about 1,000 to about 3,100 and the central hydrophic being about 10 44% to both 50 44% of the final molecule such as the PLURONIC(R) R surfactants and the TETRONIC(R) R (ethylene oxide and propylene oxide with ethyleneodiamine justractants also available from BASF Corp.; and

min cuty relocationly during state as set of earth and in the Confed by "capping" or "end blocking" the terminal hydroxy group or groups by reaction with small hydroxylobic molecules such as propylene oxide, butylene oxide, benry/chorfice, short chain latty acids, alconidos or alght haldes containing from 1 to about 5 carbon atoms, converting terminal hydroxy groups to chloride with thionyl chloride, and so forth leading to all-block, blockhetein, bettero-block or all-heteric nonlonicis.

[0051] More suitably, the nonionics useful herein include, but are not limited to, block copclyment of ethylene oxide and propylene oxide sequentially condensed upon initiators having difunctional or tetrafunctions reactive hydrogens and alcohol alkoxylates. Especially preferred surfactarts for compositions of the present invention are mixtures of alityl sulforates and block copolymens of ethylene oxide and provivere oxide sequentially condensed onto an ethylenediamine initiation.

[0052] A bend of surfactants may be suitably employed in the present invention to anne at the characteristics desirable for a particular application, For instance, some embodiments may include a surfactant for emulative application, For instance, some embodiments may include a surfactant for emulative and the property of the propert

10059] As noted above, in some embodiments of the present invention, a coupler or hydrotrope will suitably be added to the compessions, particularly when supplied in concentrated from its obtailize the stip adds in water. Those which have been found to be particularly effective for solubilizing the fatly adds of the present invention include, but are not limited, to the arrival not include and a stip adds and the stip adds of the present invention include, but are not limited, to the arrival not include such as the add and the state of the stip adds of the stip adds and the stip adds of the sti

[DOS4] Commercially available hydrotoce/couplers include, for example, DOWFAX(R) allysteed deliberate oxide disulficates surfactants; PETROL(R) silvi inphiritiene sulfanes surfactants; BIOTERGE(R) PAS-8 octane sulforate surfactants; and so forth. The proportion of the surfactant component which is made up hydrotropic depends upon various factors including the expeditic hydrotiops employed, and the specific factor acid employed; for instance. This hydrotoce should be specifically suicellar from \$% ox about \$4 \times 4 \times

[00:55] The addition of an anionic hydrotrope has been found useful for maintaining product stability, and for decreasing the chance of phase separation over time. [00:65] The lists of ingredients given herein are intended as exemplary lists and are by no means exhaustive

of the ingredients usuful herein. Such lists are not intended to limit the scope of the present invertion. [Option John Lines and the present invertion (DOST) Other ingreders may be opionally added to the compositions of the present invention to impart additional properties to the composition in amounts which do not detrimentally affect the desired properties obtain properties or properties in amounts which do not detrimentally affect the desired properties obtained by the properties of the desired properties obtained by the properties of the desired properties of the properties of the

[0.058] The balance of the concentrates and/or disted use solutions is typically valer. A concentrate may or may not comprise any water. The concentrates may be disted with any amount, but are typically disted in the range of about 1:100 to about 1:1500 parts concentrate to water which are typical of normal use indistinces. For clearing compositions, the compositions are typically more concentrations for example, clearing compositions may be distinct to ratios of about 1:100 to about 1:500, more saliably about 1:500. The clearing compositions of the compositions are typically more concentration of the clearing compositions of the compositions are typically more constantially about 1:500. The contract the cliability of the compositions are typically compositions of the composition of the composition of the composition of water (2.57\*10\*-0.2 \*) liters to about 3.785 liters or about 2.5.57 m to about 3.785,41 m). This ratio is approximately 1.788 parts.

is the most commonly used solvent for dilution.

[0059] The compositions of the present invention may be prepared in various forms in both ready-to-use, and in concentrated versions. As noted above, the concentrated compositions require no dilution, but are

typically formulated in one of several ways. (0060) Commonly, the compositions are prepared as liquid concortates intended for further mixton just prior to use, or are prepared as a prepared as the property of the prepared property of the pr

[0062] The present invention contemplates methods of using the composition for hard surface cleaning and/or sanitizing of in-place or clean-in-place (CIP)/SIP (steam-in-place) assemblies. The compositions may

be introduced into a dealing and/or sanitizing system either manually, or using an automatic metering and/or dispensing system. The compositions may be either pre- or post-disked with water before or after addition to the system. This is usually accomplished at ambient temperatures. The composition is then circulated through the system, dained, and optionsity, the system is made one or more times with potable compositions may be employed where fearing is not a concern and are contemplated as being within the soope of the present invention as described above. For example, high fooming sanitizers may be employed for sanitizing external surfaces of equipment, ceilings, walls, floors, and so forth, while low fearing compositions may be employed for cleaning the infirmal equipment systems such as piging systems, i.e.,

[0063] The present invention also contemplates methods of using the compositions as one-step classification and disinfectants in which one composition can both clean and smitize a surface simultaneously. Typically the surface is characterized as a hard surface. Such surfaces include equipment involved in both food and beverage processing such as in dairy operations including pipelines and bulk tanks and brownins.

tains and occurred and the control of the control o

[0065] The following non-limiting examples further illustrate the present invention.

Test Methods

[0066] 1. Foaming Evaluation

[00x7] Distilled water (300 m.l.) at a temperature of about 50-70(leg.) If, was poured into a 500 mt. organizated cylinder. Provideer provider (10 g.) or liquid product (10 mt.) was poured into the graduated cylinder which was then stoppered lightly. The cylinder was then inverted and returned to an upright position 10 times. The graduated cylinder was then allowed to sit and the water and form layers allowed to separate. The height of the foam layer in mt. was determined at the highest and lowest points after the designated elapsed time. The average of the two readings was reported.

[0068] foam height=(mL of foam+liquid)-(mL of liquid)

[008] 2. Food Certine Ortaan Surface Santitizing Efficacy at 25[deg.] F.
[0070] Testing was conducted according to AOAC Germicidal and Detergent Santitizing Action of
Disinfectants 950.09, Official Methods of Analysis of the AOAC International, 16thEdition, 1995. Testing

required EPA Pesticide Assessment Guidelines, Subdivision G 91-2(k)(2).

[0071] All of the examples tested were diluted at a ratio of 1 oz to 6 gallons concentrate to water (0.13%) using 500 pm synthetic hard water (as CaCo). The hard water was prepared as follows:

Hard Water Preparation

500 ppm synthetic hard water (as CaCO3) PREPARATION

Total Hardness

Solution A Solution B Volume Determination

(mL) (mL) (mL) Final pH (ppm CaCO3)

5 4 1000 7.91 500 ppm

[0072] Solution A: [0073] 31.74 g MgCl2 (or equivalent of hydrates)+73.99 g CaCl2 (or equivalent of hydrates) and diluted to 1 lifter in boiled deionized water (heat sterilized)

[0074] Solution B: [0075] 56,03 g NaHCO3 diluted to 1 liter in boiled delonized water (filter sterilized)

[0076] Two test systems were used for this study per USEPA Pesticide Assessment Guidelines Subdivision G, Series 91, Subseries 91-A, 91-2, (k)(2). FMII 1

[007] Each of the following compositions was prepared by admixing listed chemicals in sequential order, blending throughly by agritation and allowing each imprient to completely disperse or dissolve into fliquid mixture before addition of the next ingredient. The resultant compositions were clear and homogeneously uniform upon admixture of all listed impredients. The concentrates were conditioned at 40(seq.) F. until visual phase instability was observed or after 4 days with no visual change in stability. Use districts were prepared similarly using it conflicts the concentrate per organized visually probability to the concentrate per special probability. The concentrate per special probability to the concentrate per special probability to the concentrate per special probability. The concentrate per special probability of the stability results obtainable with compositions of the present invention. Variation vise excitated in the range, however, particularly in relation to the amount of time a composition remained at low temperatures. Stabilities varied and lesser stabilities were obtained depending on conditions, time, and composition in composition.

[0078] The following table 1 illustrates compositions of the present invention which utilize nitric acid as the strong acid and which contain no phosphoric acid.

TABLE 1

Glacial acetic acid 15 15 15

Nitric acid, 42[deg.] Be 30 30 30

1-octane sulfonate, sodium, 25 25 25

40% active

Perlamonic acid (C9) 6 6 6

```
Decanoic acid (C10) 1 1 1
Urea - 0.1 -
FD&C yellow #5, 0.10% - 0.44 -
(dye)
DI water 23.00 22.46 23
Comment CDS dye not stable CDS
CDS = concentrate and dilution stable, no visible precipitate/floc, very minor surface
Examples 4-8
[0079] The following table 2 illustrates compositions of the present invention that utilize a blend of nitric acid
and phosphoric acid and which contain no urea.
  TABLE 2
  45678
 Phosphoric acid, 75% 15 15 10 10 15
 Nitric acid. 42 deg Be 15 15 21 21 15
 Glacial acetic acid 15 15 15 15 15
 1-octane sulfonate, 25 25 25 25 25
 sodium, 40% active
 Perlargonic acid (C9) 6 6 6 7 7
 Decanoic acid (C10) 1 1 1 1 1
 FD&C Yellow #5, 0.10% 0.44 - - - -
 Di water 22.56 23 22 21 22
 Comments dye not CDS CDS CDS CDS
  stable
 CDS = concentrate and dilution stable, no visible precipitate/floc, very minor surface
Examples 9-18
[0080] The following table 3 illustrates compositions of the present invention having a nitric acid/phosphoric
acid blend and containing varying amounts of urea.
  TABLE 3
  9 10 11 12 13 14 15 16 17 18
 Phosphoric 10 10 10 10 10 10 10 10 10 10
 acid, 75%
 Nitric acid, 21 21 21 21 21 21 21 21 21 21
 42 deg Be
 Glacial 15 15 15 15 15 15 15 20 20 20
 acatic acid
 1-octane 25 25 25 25 25 30 25 25 30 30
 sulfonate.
 sodium, 40%
 active
 Perlargonic 7 7 7 7 7 7 6 7 7 6
 acid (C9)
 Decanoic 1 1 1 1 1 1 1 1 1 1
 acid (C10)
 Urea 0.10 0.50 1.00 5.00 0.25 0.5 0.5 0.5 0.5 0.5
 DI water 20.9 20.5 20.0 16.0 20.75 15.50 21.50 15.50 10.50 11.50
```

```
CDS = concentrate and dilution stable, no visible precipitate/floc, very minor surface
Examples 19-28
[0081] The following tables 4-8 illustrate compositions of the present invention employing various surfactant
blends
  TABLE 4
  19 20 21 22 23 24 25
 Phosphoric acid. 75% 10 10 10 10 10 10 10
 Nitric acid. 42 deg Be 21 21 21 21 21 21 21 21
 Glacial acetic acid 15 15 15 15 15 15 15
 1-octane sulfonate. 25 25 25 25 25 25 25
 sodium, 40% active
 PLURAFAC (R) RA-40 1.00 - - - - -
 TETRONIC (R) 1307 - 2.00 - - - - -
 TETRONIC (R) 1107 - - 2.00 1.00 0.50 - -
 TETRONIC (R) 908 - - - - 0.50 0.65
 Perlargonic acid (C9) 7 7 7 7 7 7 7
 Decanoic acid (C10) 1 1 1 1 1 1 1
 Urea 0.5 0.5 0.5 0.5 0.5 0.5 0.5
 DI water 19.50 18.50 18.50 19.50 20.00 20.00 19.85
 Comments ECDS ECDS ECDS ECDS ECDS ECDS
 ECDS = excellent concentrate and dilution stability, no visible precipitate/floc, no visible surface oiling at
40[deg.] F.
[0082] PLURAFAC(R) RA-40 is an alcohol ethoxylate.
[0083] TETRONIC(R) 908, 1107 AND 1307 are all nonionic surfactants block copolymer adducts of
ethylene oxide and propylene oxide to ethylenediamine.
Example 24 and Comparative Examples A-C
[0084] Comparative Examples A-C are representative of commercially available sanitizing compositions
which are standards in the industry.
 TABLE 5
 Comparative A (wt-%) Comparative B Comparative C
 30% phosphoric acid 16% soft water 11% soft water
 21,99956% citric acid, 38% phosphoric acid, 35% phosphoric
    acid
 (50% active) (75% active) (75% active)
 9% citric acid, anhydrous 10% propionic acid 8% lactic acid, food
    grade (88% active)
 30% 1-octane sulfonate, 3% perlargonic acid 34% sodium linear
  sodium (40% active) alkvl naphthalene
    sulfonate
  6% octanoic acid 3% decanoic acid 9% octanoic acid
 2% decanoic acid 30% 1-octane sulfonate, 3% decanoic acid
   sodium (40% active)
 q.s. isopropyl alcohol q.s. FD&C yellow #5
  q.s. FD&C yellow #5
[0085] A foaming evaluation was conducted according to Test Method #1. FIG. 1 is a bar graph showing the
 results of the foaming evaluation. As can be seen from the graph, example 24 exhibited a lower foam height
```

100861 Formula 24 was further tested for food contact surface sanitizing efficacy at 25[deg.] F. as described

TABLE 6

than comparatives A-C which are standards in the industry.

in Test Method #2 above. The following results were obtained.

```
Efficacy Test Results
```

Staphylococcus aureus ATCC 6538

Test Substance Average CFU/mL of Test

(Batch Number) Survivors Percent Reduction

Formula 24 3.0 \* 10<1> >99.999

(Batch 1) 1.8 \* 10<2> >99.999

Formula 24 7.5 \* 10<1> >99.999

(Batch 2) 7.5 \* 10<1> >99,999

Formula 24 5.5 \* 10<1> >99.999

(Batch 3) 2.8 \* 10<2> >99.999

TABLE 7

Efficacy Test Results

Escherichia coli ATCC 11229

Test Substance Average CFU/mi\_ of Test

(Batch Number) Survivors Percent Reduction

Formula 24 7.5 \* 10<1> >99,999

(Batch 1) <10 >99.999

Formula 24 <10 >99,999

(Batch 2) <10 >99.999

Formula 24 <10 >99.999

(Batch 3) <10 >99.999

[0087] As can be seen from tables 6 and 7, formula 24 exhibited a 99.999% reduction of S. aureus and E. coli. Example 24 therefore meets the efficacy requirements of a food contact surface sanitizer.

TABLE 8

26 27 28

Phosphoric acid, 75% 10 10 10

Nitric acid, 42 deg Be 21 21 21

Gladal acetic acid 15 15 15

1-octane sulfonate, sodium, 25 25 25

40% active

DOWFAX (R) C-6 add, - - -

45%

DOWFAX (R) C-6 - - -

\_\_\_\_\_

acid, 78%

PETRO (R) LBA liquid, 50% - - -

TETRONIC (R) 908 1.00 1.50 2.00

Perlargonic acid (C9) 7 7 7

Decanoic acid (C10) 1 1 1

Urea 0.5 0.5 0.5

DI water 19.50 19.00 18.50

Comments ECDS ECDS ECDS

DOWFAX (R) C-6 is a sodium hexyl diphenyloxide disulfonate

PETRO LBA is a sodium alky naphthalene sulfonate

TETRONIC (R) 908 is a block copolymer adduct of ethylene oxide and propylene oxide to ethylenediamine.

TABLE 1

Cleaning Compositions

Example 29 Example 30

Phosphoric acid, 75% 20.0 20.0

Nitric Acid, 42 Be 21.0 21.0

Glacial Acetic Acid 15.0 15.0

1-octane sulfonate 10.0 20.0

TETRONIC (R) 908 0.5 1.50

Perlargonic acid 1.0 3.4

Decanoic acid 0.15 0.5

Urea 0.5 0.5

DI water 26.85 18.10 [0.088] The above compositions are illustrative of compositions of the present invention which are useful as cleaning compositions, i.e. one-step cleaning compositions. Example 29 is intended for 1% dilution (1:100 concentrate to water) and example 29 is intended for 0.3% (1:333 concentrate to water) dilution.

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# Acid sanitizing and cleaning compositions containing protonated carboxylic acids

The EPO does not accept any responsibility for the accuracy of data and information originating from other authorities than the EPO, in particular, the EPO does not guarantee that they are complete, up-to-date or fit for specific purposes.

Claims not available for JP 2005511635 (T) Claims of corresponding document: US 6472358 (B1)

# What is claimed is:

[0089] 1. A sanitizing and/or cleaning composition comprising a) about 3 wt% to about 12 wt% of the composition of at least one alignate short clean antimicrobally effective C5 to C14 fathy acid or mixture three(f) b) about 5 wt% to about 50 wt% of the composition of at least one walk C1 to C4 carboxylic acid; and c) about 5 wt% to about 5 wt% of the composition of a strong acid which is nitric acid or a mixture of nitric and phosphoric acids.

[0090] 2. The composition of claim 1 wherein said at least one weak carboxylic acid is acetic acid, hydroxyacetic acid, projionic acid, hydroxyacetic acid, acid,

[0091] 3. The composition of claim 1 wherein said at least one aliphatic short chain antimicrobially effective fatty acid is a C6 to C10 fatty acid.

[0092] 4. The composition of claim 1 wherein said at least one weak carboxylic acid is acetic acid.

[0093] 5. The composition of claim 1 wherein said aliphatic short chain fatty acid is decanoic, nonanoic or a mixture thereof.

[0094] 6. The composition of claim 1 wherein said aliphatic short chain fatty acid is nonanoic acid.

[0095] 7. The composition of claim 1 wherein said aliphatic short chain fatty acid is present at a concentration of about 5 wt-% to about 10 wt-% of the composition.

[0096] 8. The composition of claim 5 wherein said mixture comprises about 0.25 to about 5 wt-% of the composition decanoic acid and about 2 to about 10 wt-% of the composition nonancic acid.

[0097] 9. The composition of cialm 5 wherein said mixture comprises about 0.5 to about 4 wt-% of the composition decanoic acid and about 3 to about 9 wt-% of the composition nonanoic acid.

[0098] 10. The composition of claim 5 wherein said mixture comprises about 1 to about 3 wt-% of the composition decanoic acid and about 4 to about 8 wt-% of the composition nonanoic acid.

[0099] 11. The composition of claim 5 wherein said mixture comprises about 1 wt-% of the composition decanoic and about 6 to about 7 wt-% of the composition nonanoic acid.

[0100] 12. The composition of claim 1 wherein said at least one weak carboxylic acid is present at a concentration of about 10 wt-% to about 40 wt-% of the composition.

I0101113. The composition of claim 1 wherein said at least one weak carboxylic acid is acetic acid.

[0102] 14. The composition of claim 1 wherein said phosphoric acid is present at a concentration of about 5 wt-% to about 40 wt-% of the composition.

[0103] 15. The composition of claim 1 wherein said phosphoric acid is present at a concentration of about 10 wt-% to about 35 wt-% of the composition.

[0104] 16. The composition of claim 1 wherein said nitric acid is present at a concentration of about 5 wt-% to about 50 wt-% of the composition.

[0105] 17. The composition of claim 1 wherein said nitric acid is present at a concentration of about 15 wt-% to about 40 wt-% of the composition.

[0106] 18. The composition of claim 1 further comprising an effective amount of urea to reduce nitrogen peroxide to nitrogen.

[0107] 19. The composition of claim 18 wherein said effective amount of urea is about 0.05 wt-% to about 5 wt-% of the composition.

[0108] 20. The composition of claim 18 wherein said effective amount of urea is about 0.5 wt-% of the

[0109] 21. The composition of claim 1 further comprising at least one surfactant.

[0110] 22. The composition of claim 21 wherein said at least one surfactant is nonionic

[0111] 23. The composition of claim 22 wherein said at least one surfactant is a tetrafunctional block copolymer derived from the addition of ethylene oxide and propylene oxide to ethylenediamine.

[0112] 24. The composition of claim 21 wherein said surfactant is present at a concentration of 0.1 wt-% to about 50 wt-% of the composition.

[0113] 25. The composition of claim 1 further comprising at least one organic hydrotrope.

- [0114] 26. The composition of claim 25 wherein said organic hydrotrope is an anionic sulfonate or corresponding disulfonate.
- [0115] 27. The composition of claim 26 wherein said organic hydrotrope is an allyl sulfonate, an any sulfonate, a C6-30 allaryl sulfonate or a corresponding disulfonate, an allylated diphenyl oxide disulfonate, or an anionic mono or disubstituted allyl ethoxylated phosphate ester, or a mixture thereof.
- [0116] 28. The composition of claim 27 wherein said organic hydrotrope is 1-octane sulfonate.
- [0117] 29. The composition of claim 25 wherein said organic hydrotrope is present at a concentration of about 0.5 wt-% to about 50 wt-% of the composition.
- [0118] 30. The composition of claim 1 further comprising water.
- [0119] 31. The composition of claim 30 wherein said concentrate is diluted with water at a ratio of 1:100 to about 1:1500 parts concentrate to water.
- [0120] 32. The composition of claim 30 wherein said concentrate is diluted at a ration of about 1:768 parts concentrate to water.
- [0121] 33. A method of one-step cleaning and sanitizing a surface comprising the step of contacting said surface with a composition as in claim 1.
- [0122] 34. The method of claim 33 further comprising the step of diluting said composition with water at a ration of about 1:100 to about 1:1500 of the composition to water.
- [0123] 35. A clean-in-place method of cleaning a bewerage or food processing unit including conduits, surfaces and containers, comprising the steps of any providing the composition of claim 1; and b) contacting conduits, surfaces and containers in said bewerage processing unit, c) removing said composition from said unit for the purpose of reintilitating processing.
- [0124] 36. The method of claim 35 further comprising the step of diluting said composition with water at a ratio of about 1:100 to about 1:1500 of the composition to water.
- [0125] 37. A distrable acid santifizing and/or deaning concentrate composition comprisingations 0.52 ke-% to about 10 km% of the concentrate of at least one C6 to C10 fatly acid or mixture thereof, about 5 km% of bacut 50 km% of the concentrate of a C1 to C4 week carboxylic acid, about 0 km% or about 40 km% of the concentrate principhoric acid; about 5 km% of the concentrate principhoric acid; about 5 km% of the concentrate principhoric acid; and about 0.05 km% to about 5 km% of the concentrate urea;
- [0126] with the proviso that the concentration of nitric acid and phosphoric acid does not exceed about 50 wt-% of the concentrate.
- [0127] 38. The composition of claim 37 wherein said phosphoric acid is present at a concentration of about 5 wt-% to about 40 wt-% of the concentrate.
- [0128] 39. The composition of claim 37 further comprising at least one surfactant at a concentration of about 0.1 wt-% to about 50 wt-% of the concentrate.
- [0129] 40. The composition of claim 37 further comprising water.
- [0130] 41. A cleaning composition comprising at about 0.5 w/% to about 10 w/% of at least one short chain first actic; 0 to w/% to about 40 w/% phosphore actic; 0 about 5 w/% to about 50 w/% entire actic; 1 about 50 w/% to about 50 w/% of at least one surfactant; and e) about 5 w/.% to about 50 w/.% of the concentrate of a C1 to C4 week carboxylic actic.
- [0131] with the proviso that the concentration of nitric acid and phosphoric acid does not exceed about 50
- [0132] 42. The composition of claim 41 wherein said composition comprises about 5 wt-% to about 40 wt-% of said at least one surfactant.
- [0133] 43. The composition of claim 42 wherein said at least one surfactant is anionic.
- [0134] 44. The composition of claim 43 wherein said at least one surfactant is an alkyl sulfonate, an aryl sulfonate, a C6-30 alkaryl sulfonate or a corresponding disulfonate, an alkylated dipheryl oxide disulfonate, or an anionic mono or disubstituted alkyl ethoxydated phosphate ester, or a mixture thereof.
- [0135] 45. The composition of claim 44 wherein said at least one surfactant is a 1-octane sulfonate.
- [0136] 46. The composition of claim 41 wherein said at least one surfactant is nonionic.
- [0137] 47. The composition of claim 46 wherein said at least one surfactant is a tetrafunctional block copolymer derived from the addition of ethylene oxide and propylene oxide to ethylenediamine.
- [0138] 48. The composition of claim 46 wherein said at least one surfactant is present at a concentration of about 0.25 wt-% to about 10 wt-%.
- [0139] 49. The composition of claim 41 wherein said at least one fatty acid is present at a concentration of about 1 wt-% to about 5 wt-%.
- [0140] 50. The composition of claim 41 wherein said at least one fatty acid comprises about 0.1 wt-% to about 5 wt-% decancic acid and about 0.5 wt-% to about 10 wt-% nonanoic acid.
- [0141] 51. The composition of claim 41 wherein said at least one fatty acid comprises about 0.1 wt-% to about 1 wt-% decanoic acid and about 1 wt-% to about 5 wt-% nonanoic acid.

[0142] 52. A sanitizing and/or cleaning composition comprising:a) 1 part of at least one aliphetic short chain antimicrobially effective C5 to C14 fatly acid or mixture thereof; b) about 1.7 to about 4.2 parts of at least one weak C1 to C4 cartoxylic acid, and Q) about 1.7 to about 4.2 parts of a strong acid which is nitric acid or a mixture of nitric and phosphoric acids.

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Corresponding U.S. Petent



# (12) United States Patent Richter et al.

(51) Int. Cl.7 .....

(58) Field of Search .....

(10) Patent No.: US 6,472,358 B1 (45) Date of Patent: Oct. 29, 2002

(54)	ACID SA	NITIZING AND CLEANING	(56)				Referer	ices Cited
		ITIONS CONTAINING ATED CARBOXYLIC ACIDS			U.S	i. I	PATENT	DOCUMENTS
				,002,775				Kabara 426/532
(75)	Inventors:	Francis Lawrence Richter, Lino		,343,798				Fawzi 424/240
		Lakes, MN (US); Duane Joseph		,404,040				Wang 134/22.14
		Reinhardt, Manlewood, MN (US);		,406,884				Fawzi et al 424/81
		Richard R. Staub, Lakeville, MN		,822,513				Corby
		(US); Teresa C. Podtburg, Waconia,		,208,257				Kabara
		MN (US)		,308,868				McKinzie et al 424/605
		(,		391,379				McKinzie et al 424/605
(73)	Assignee:	EcoLab Inc., Mendota Heights, MN		462,714				Talwalker 422/37
` ′	_	(US)		,569,461				Andrews 424/405
(*)	Notice:	Subject to any disclaimer, the term of this	* cite	i by exa	mir	cr		
(*)	Notice:	patent is extended or adjusted under 35	Prima	rv Fran	ino	r-	-Charles	Boyer
		U.S.C. 154(b) by 0 days.						m-Vidas, Arrett & Steinkraus
		0.5.C. 154(6) by 0 days.	P.A.	morney,	***	٠,,	, 0, 1,	" vidus, raron de olemanaus
(21)	Appl. No.:	10/000,138	(57)				ABS:	TRACT
(22)	Filed:	Nov. 15, 2001	A san	itizing c	om	008	sition co	mprising at least one aliphatic

..... C11D 7/08

510/247; 424/718

..... 510/234, 253, 510/258, 247; 424/718

short chain antimicrobially effective  ${\rm C_3}$  to  ${\rm C_{14}}$  fatty acid or mixture thereof, at least one carboxylic weak acid and a strong mineral acid which may be nitric or a mixture of nitric and phosphoric acids.

52 Claims, 1 Drawing Sheet

30 Minute I Minute Initial Example 24 Comp C Fig. 1 Foaming Evaluation 78 Comp B 50 78 Comp' A 50 86

Foam Height mLs

# ACTO SANITIZING AND CLEANING COMPOSITIONS CONTAINING PROTONATED CARBOXYLIC ACIDS

### FIELD OF THE INVENTION

The present invention relates to acid sanitizing and/or cleaning compositions comprising antimicrobially effective Cs to C4 carboxylic acids. The present invention is directed to both concentrates and to water diluted use solutions.

### BACKGROUND OF THE INVENTION

Periodic cleaning and sanitizing in food, drink, pharmaceutical, cosmetic and similar processing industries; in food preparation and service businesses; in health and day care facilities; and, in hospitality establishments are a necessary practice for product quality and public health. Residuals left on equipment surfaces or contaminants found in the process or service environment can harbor and nourish growth of subsequent processed product or critical contact 20 surfaces. Protecting the consumer against potential health hazards associated with pathogens or toxins and maintaining the quality of the product or service requires routine clean-ing of residuals from surfaces and effective sanitation to reduce microbial populations.

Visual inspection of the equipment cannot ensure that surfaces are clean or free of microorganisms. Antimicrobial treatments as well as cleaning treatments are therefore required for all critical surfaces in order to reduce microbial population to safe levels established by public health regu-lations. This process is generally referred to as sanitizing. The practice is of sanitation is particularly of concern in food process facilities wherein the cleaning treatment is followed by an antimicrobial treatment applied upon all critical surfaces and environmental surfaces to reduce the microbial population to safe levels established by ordinance. A sanifized surface is, as defined by the Environmental Protection Agency (EPA), a consequence of a process or program containing both an initial cleaning and a subsequent samtizing treatment which must be separated by a potable water rinse. A sanitizing treatment applied to a cleaned food 40 contact surface must result in a reduction in population of at least 99.999% (5 log) for specified microorganisms as defined by the "Germicidal and Detergent Sanitizing Action of Disinfectants", Official Methods of Analysis of the Association of Official Analytical Chemists, paragraph 960.09 45 matrix which has the potential of harboring bacteria, an and applicable sections, 15th Edition, 1990 (EPA Guideline effect opposite to that desired. 91-1).

The antimicrobial efficacy of sanitizing treatments is significantly reduced if the surface is not absolutely free of soil and other contaminants prior to the sanitizing step. The 50 presence of residual food soil and/or mineral deposits inhibit sanitizing treatments by acting as physical barriers which shield microorganisms lying within the organic or inorganic layer from the microbicide. Furthermore, chemical interactions between the microbicide and certain contaminants can disrupt the killing mechanism of the microbicide.

With the advent of automated clean-in-place and sanitizein-place systems, the need for disassembly has been diminished, and cleaning and sanitizing have become much more effective. However, modern food industries still rely on sanitizers to compensate for design deficiencies or operational limitations in their cleaning programs and the probability of very small residual amounts of organic and inorganic soils and biofilms remaining on food contact surfaces after cleaning. In cooperation with these process changes and higher performance expectations, sanitizer treatments 65 must also comply with the increasing demand for safer, less corrosive, more environmentally friendly compositions.

According to the U.S. Center for Disease Control and Prevention, food poisoning in calendar year 2000 resulted in 5000 deaths, 325,000 hospitalizations and 76,000,000 illnesses. The need exists for improved sanitizing treatments to destroy pathogens and food spoilage microorganisms resistant to conventional treatments within the food gathering, food processing, and food serving industries. A further complication is that the list of approved microbicidal agents has continued to decrease due to acute and chronic human toxicity of some microbicidal agents, and to their environmental persistence in water supplies.

Antimicrobially active acids have been used in sanitizing operations, For instance, U.S. Pat. No 404,040 describes a operations. For instance, U.S. Pat. No 404,040 describes a sanitizing composition comprising aliphatic, short chain fatty acids, a hydrotrope or solubilizer for the fatty acids, and a hydrotrope-compatible acid, and U.S. Pat. No. 5,330,769 describes fatty acid sanitizer concentrates and diluted final solutions which include individual amounts of germicidally effective fatty acid, hydrotrope, a strong acid group consist-ing of phosphoric acid and sulfuric acid or mixtures thereof sufficient to lower the pH of the final solutions to about 1-5, and a concentrate stabilizing weak acid component selected from the group consisting of propionic, butyric and valeric acids and mixtures thereof.

Protonated carboxylic acids offer broad spectrum antimicrobial activity against gram-positive and gram negative bacteria, persistent biocidal activity in the presence of organic and inorganic soils and residual biocidal and inhibitory activity. They combine both acid for mineral deposit control and sanitizer for antimicrobial effect into one treatment solution

However, one problem associated with the use of protonated carboxylic acid sanitizers is poor use dilution phase stability, particularly at lower water temperatures of 40°-50° F. 50° F. Fatty monocarboxylic acids having alkyl chains containing 5 or more carbon atoms, are typically characterized as water insoluble and can oil out or precipitate from solution as a gelatinous flocculent. Solubility tends to decrease with decreasing water temperature and increasing ionic concentration. Furthermore, the oil or precipitate can affix to the very surfaces which the sanitizing solution is intended to sanitize, such as equipment surfaces, leading to a film formation on these surfaces over time. The fatty acid film deposited and left remaining on the equipment surface tends to have a higher pH than the sanitizing solution from which it came resulting in a significantly lowered biocidal efficacy, and, if mixed with food soil, may result in a film

One solution has been to use short chain, C1-C4 carboxylic or hydroxycarboxylic acids to solubilize and thus stabilize longer chain fatty acids in high actives composition concentrates. However, these short chain weak acids have been known to be less effective at normal use dilution concentrations than their longer chain counterparts, and extreme dilution of the concentrate in water leads to a decrease in the solvating effect resulting in a precipitate of the longer chain length fatty acids of Cs or higher from solution. Furthermore, raising the concentration of the C<sub>1</sub>-C<sub>4</sub> acids increases the cost of the sanitizing composition, and does not appear to result in a significant increase in dilution stability or to improved antimicrobial efficacy.

Organic hydrotropes or coupling agents, such as low molecular weight sulfonates, may be employed to increase the solubility and miscibility of the longer chain fatty acids with water and inorganic salts both in concentrated and in diluted use solutions. Again, the solubility appears to diminish at sustained lower water temperatures with the result being phase separation.

There remains a need in the art for an improved biocidal composition which utilizes a carboxylic fatty acid which has 3

high antimicrobial efficacy, has good phase stability, exhibits low toxicity, and is not detrimental to the environment.

# SUMMARY OF THE INVENTION

Surprisingly, the compositions of the present invention < exhibit excellent phase stability both in concentrated form and in diluted use solutions, and in particular, they exhibit excellent phase stability in low temperature water diluted use solutions. Even more surprisingly, the stability is improved in the presence of nitric acid.

The sanitizing and/or cleaning compositions of the present invention, in both concentrated and in diluted use solutions, include an antimicrobially effective short chain fatty acid, a shorter chain weak carboxylic acid, and a strong mineral acid. The shorter chain weak carboxylic acid functions as a solvent.

The shorter chain weak carboxylic acid functions as a solvent for the antimicrobial short chain fatty acid. In concentrated form, the compositions also desirably contain an organic hydrotrope.

In some embodiments, the antimicrobially effective short  $^{20}$ chain fatty acid is a C5 to C14 fatty acid, and more suitably C6 to C10 fatty acid, or some mixture thereof, the shorter chain weak carboxylic acid is a C, to C, carboxylic acid, and the strong mineral acid is nitric, or a mixture of nitric and phosphoric acids.

In some embodiments wherein a hydrotrope is included in the composition, an anionic sulfonate hydrotrope is employed.

Additionally, the composition may optionally include at least one anionic and/or nonionic surfactant. In some 30 embodiments, a nonionic surfactant is suitably employed to improve surface wetting, soil removal, and so forth. It may also function to improve the solubility of the fatty acids at

The antimicrobailly effective effective short chain fatty 35 acid is useful from about 3 wt-% to about 12 wt-% of the concentrate, and more suitably from about 5 wt-% to about 10 wt-% of the concentrate. In one particular embodiment, the concentrate includes a blend of two fatty acids.

The weak carboxylic acid is useful from about 5 wt-% to 40 about 50 wt-% of the concentrate, and more suitably from about 10 wt-% to about 40 wt-% of the concentrate. In one particular embodiment, the weak carboxylic acid component includes at least acetic acid. The weak carboxylic acid acts

The strong mineral acid is useful from about 5 wt-% to about 50 wt-% of the concentrate, and more suitably about 15 wt-% to about 40 wt-% of the concentrate. In some embodiments, the strong mineral acid is nitric which is useful from about 5 wt-% to about 50 wt-% of the concentrate, and more suita concentrate. If phosphoric acid is employed, it is useful from 5 wt-% to about 40 wt-% of the concentrate, and more suitably about 10 wt-% to about 35 wt-% of the concentrate

Surprisingly, the antimicrobially active short chain fatty 55 acid is stable in nitric acid

The compositions may further comprise optional ingredients including urea for stabilization of nitric acid, and a surfactant component. The surfactant component may include one or more surfactants. In some embodiments, an 60 anionic or nonionic surfactant may be optionally added at a level of 0.1 wt-% to about 50 wt-% of the concentrate, more suitably about 0.25 wt-% to about 40 wt-% of the concentrate, even more suitably about 0.5 wt-% to about 40 wt-%, and most suitably about 1 wt-% to about 30 wt-%.

In some embodiments, an anionic hydrotrope is employed at a level of about 0.5 wt-% to about 50 wt-%, suitably about

1 wt-% to about 40 wt-% of the concentrate, and more suitably from about 5 wt-% to about 30 wt-% of the concentrate. In one embodiment, the anionic hydrotrone includes at least one alkylsulfonate.

The compositions may be diluted with water at any ratio whatsoever, but typically the ratio is between about 1:100 parts of the concentrate to water to about 1:1500 parts of the concentrate to water. This is referred to as a use dilution. A very typical use dilution is about 1 ounce of concentrate to about 6 gallons of water which is a ratio of about 1:768 parts

of the concentrate to water

The compositions of the present invention find utility as both sanitizing and disinfecting compositions as well as cleaning compositions, and are useful for both hard and soft surface sanitizing and disinfecting in farm operations, food processing operations, institutional food preparation and serving areas, health care and child care facilities as well as any other number of contact sensitive environments. The compositions exhibit high antimicrobial efficacy while having low toxicity, are not detrimental to the environment, and do not contaminate food stuffs.

The compositions also find utility for use as one-step cleaning/sanitizing compositions and disinfectants in which the composition cleans and sanitizes simultaneously.

## BRIEF DESCRIPTION OF THE FIGURES

FIG. 1 is a bar graph illustrating results of a foaming evaluation conducted for example 24 which was compared to three commercially available sanitizing compositions.

### DETAILED DESCRIPTIONS OF THE PREFERRED EMBODIMENTS

While this invention may be embodied in many different forms, there are described in detail herein specific embodiments of the invention. This description is an exemplification of the principles of the invention and is not intended to limit the invention to the particular embodiments illustrated.

The antimicrobial agents useful herein include those referred to generally in the art as acid-anionics including carboxylic acids having biocidal activity when in protonated form. These antimicrobial agents are typically classified as having low toxicity and as being environmentally friendly.

The term "short chain fatty acids" as used herein refer to those acids generally having about 5 to 14 carbon atoms, as a solvent for the antimicrobially active short chain fatty 45 suitably about 6 to 12 carbon atoms, more suitably from about 6 to 10 carbon atoms and most suitably about 7-10 carbon atoms. In some embodiments of the present invention, a blend of a Co, fatty acid and a C10 fatty acid or a blend of nonanoic and decanoic acids is employed. The short chain fatty acids are useful from about 3 wt-%

to about 12 wt-% of the concentrate and suitably about 5 wt-% to about 10 wt-% of the concentrate. In some embodiments of the present invention which employ a blend, nonanoic acid is utilized from about 2 wt-% to about 10 wt-% of the concentrate, suitably from about 3 wt-% to about 9 wt-%, and more suitably from about 4 wt-% to about 8 wt-% of the concentrate while decanoic acid is utilized from about 0.25 wt-% to about 5 wt-%, suitably from about 0.5 wt-% to about 4 wt-% and more suitably about 1 wt-% 3 wt-% of the concentrate. The present inventors have found that when employing such a blend of nonanoic and decanoic acids, the phase stability appears to be improved when employing more nonanoic acid, and less decanoic acid. It is surmised that the shorter chain of the nonanoic acid provides increased solubility in water over the decanoic acid, while the decanoic acid provides increased antimicrobial efficacy over the nonanoic acid. Blending the two has been found to be particularly advantageous.

5

The carboxylic weak acid is a C, to C, carboxylic acid. Examples of suitable carboxylic weak acids include, but are not limited to, accite acid, hydroxyacetic acid, propionic acid, alpha selotpropionic acid, chias hestopropionic acid, alpha selotpropionic acid, alpha selotpropionic acid, alpha selotpropionic acid, alpha selotpropionic acid, and acid acid acid acid acid acid solvent includes accid acid. As noted above, the carboxylic weak acid acid acid solvent includes accid acid. As noted above, the carboxylic weak acid is useful years and acid acid. As noted above, the carboxylic weak acid is useful acid. The carboxylic weak acid is useful acid. The carboxylic weak acid is useful acid.

The strong acid component of the compositions is utilized to lower the pH in the final solutions to a desirable level of about 1-5, and preferably from about 2.5-4. The strong acid is suitably either nitric acid, or a mixture of nitric and phosphoric acids. Nitric acid is useful from about 5 wt-% to about 50 wt-% of the concentrate, and suitably from about 15 wt-% to about 40 wt-%. Phosphoric acid is useful from about 0 wt-% to about 40 wt-% of the concentrate and more suitably about 5 wt-% to about 35 wt-% of the concentrate. The fatty carboxylic acids of the present invention have been 20 found to be particularly stable in the presence of nitric acid due to increased solubility in the presence of nitric acid. Nitric acid is also advantageously used in the compositions of the present invention because it is economical, and because it offers added protection to stainless steel by 25 maintenance of the passive surface layer. Stainless steel is corrosion resistant due to an oxide film laver on the surface resulting from treatment with strong oxidizing agents such as nitric acid. Surfaces with this property are referred to as passive, or have a lower degree of chemical activity.

Another problem often associated with the use of acid santizers is corrosion of stainless steel surfaces associated with high mineral waters or softened waters containing chlorides which promote and accelerate corrosion of such surfaces. Strong oxidizing acids with oxidizing potential sufficient to passivate stainless are capable of reducing or eliminating such a problem.

A small amount of uses may be optionally employed in the compositions of the present invention. Organic degradation can occur in the presence of nitric acid by oxidation and diration mechanisms due to the presence and oxidizing 40 power of nitrogen dioxide (NO<sub>2</sub>) and nitrogen tetroxide (NO<sub>2</sub>), collectively referred to a sintegon peroxide. Urea may be added to react with the nitrogen peroxide to nitrogen, the nitrogen peroxide to nitrogen organically the nitrogen peroxide to nitrogen, and the nitrogen peroxide to nitrogen peroxide to nitrogen, and the nitrogen peroxide to nitrogen peroxide to nitrogen, and nitrogen peroxide to nitrogen peroxide to nitrogen, and nitrogen peroxide to nitrogen peroxide to nitrogen peroxide to nitrogen, and nitrogen peroxide to nitrogen peroxide to

Surfactants may also be optionally added to the compositions of the present invention for a variety of reasons so including improved surface wetting by lowering the surface tension, improved soil or biofilm penetration, improved soil or biofilm penetration, removal and suspension of organic soils, enhancement of biocidal effect, characterization of foam profile, i.e. by the addition of low foaming and high foaming surfactants, and increasing the solubility of the fatty acid antimicrobial in water by acting as a hydrotrope or coupling agent for the fatty acid antimicrobial to mention a few. One skilled in the art will understand that some surfactants or mixtures of surfactants serve one or more of these purposes better than others. The surfactant or mixture of surfactants selected will therefore impart different beneficial characteristics to the compositions depending on the selection made. The surfactants may be selected depending on the expected use, method of application, concentration, temperature, foam control, soil type, and so forth. The 65 selection will of course also depend on the end use application of the composition.

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The surfactants useful herein include nonionic, anionic and cationic surfactants. Most suitably, the surfactants employed include water soluble or water dispersible anionic or nonionic surfactants, or some combination thereof.

Useful anionic surfactants include, but are not limited to, those compounds having an hydrophobic group of  $C_{\rm q,20}$  such as alkyl, alkylaryl, alkenyl, acyl, long chain hydroxyalkyl, alkoyylated derivatives thereof and so forth, and at least one water-solubilizing group of acid or salt form derived from sufficine acid, ester, phosphoric acid ester and carboxylic acid. The salt may be selected based on the specific formulation to which it is being added.

More suitably, the anionic surfactants useful herein include, but are not limited to, sulfonated anionics such asklyd sulfonates or disulfonates, alkyl aryl sulfonates, alkyl anjohalene sulfonates, alkyl diphenyl oxide disulfonates, alkyl diphenyl oxide disulfonates,

More particularly, the anionic surfactants more suitable for use herein include, but are not limited to, those anionic surfactants which are linear or branched C<sub>c</sub>-C<sub>c</sub>, allylben-zue sufforates, allyl nuphtalen sufforates, long chain alkene sufforates, long chain hydroxyalkane sufforates, long chain hydroxyalkane sufforates, alkene sufforates and the corresponding disulforates including 1-octane sufforate and 12-octane disulforate; allyl suffates, allyl poly(tetyleneoxy) estifates and an aromatic special polytoxy of the suffates or condensation products of ethylene oxide and nonly phenol, having 1 to 6 oxyethylene groups per molecule, other sufforated surfactants, and so forth.

Specific examples of anionic surfactants mithible for use Specific examples of anionic surfactants withible for use berein incibie aliqy isufficates such as 1-cotane audionate to consider the surface of the surface of incibiding Stepas Co. in Northfield, Ill. under the radename of BlO-TERGE® PAS-8; PILOT® L-45, a C<sub>11,3</sub> alkylbenzene sulfonate (referred to as "LAS") from Pilot Chemical Co; BlO-SOFT® \$100 and \$130, non-neutralized linear alkylbenzene sulfonic suick (neirred to as "ILLAS"), and \$40, also an LAS, all from Stepan Company, DOVEAX® anionic variable from Dove Chemical Co, including C-6 (45% and 78%), C.—C., alkyl naphthalene sulfonates such as those variable from PetroChemicals Co. under the tradename of

PETRO® including the liquid PETRO® LBA; and so forth.

Examples of nonionic surfactants useful in the compositions of the present invention include, but are not limited to, the following classes:

 polyocxpropylene-polyocxylethylene block polymers including those made from propoxylation and/or ethoxylation of an initiator hydrogen compound such as propylene glycol, ethylene glycol, glycerol, trimethylolpropane, ethylenediamine, and so forth such as those sold under the tradename of PLURONIC® AND TETRONIC® available from BASF Core:

2) condensation products of one mole of C<sub>8</sub> to C<sub>18</sub> branched or straight chain alkyl or dialkyl phenol with about 3 to about 50 moles of ethylene oxide such as those sold under the tradename of IGEPAL® available from Rhone-Poulenc and TRITON® available from Union Carbide.

3) condensation products of one mole of a saturated or unsaturated, branched or straight C<sub>c</sub> to C<sub>2x</sub> alcobols with about 5 to about 50 moles of ethylene oxide such as those sold under the tradename of NEODOL® available from Shell Chemical Co. and ALFONIC® available from Condea Vista Co.;

4) condensation products of one mole of saturated or unsaturated, branched or straight chain C<sub>B</sub> to C<sub>18</sub> carboxylic acids with about 6 to about 50 moles of ,

ethylene oxide such as those available under the tradename of NOPALCOL® from Henkel Corp. and LIPOPEG® from Lipo Chemicals, Inc.; and other alkanoic esters formed by condensation of carboxylic acids with glycerides, glycerin, and polyhydric alcobols:

- 5) surfactants produced by the sequential addition of chylene exide and propytees coxide to ethylene glycol, ethylenediamine which result in a hydrophile with hydrophothic blocks (ac. propytees oxide) at the terminal ends (the hydrophile and hydrophothic blocks are reversed) of each molecule weighing from about 1,000 to about 3,100 and the central hydrophile being about 10 www. 4% to about 50 www. 5% of final molecules such as the PLURONIC@ Restrictants and the TEIRONIC@ 15 (Ethylene oxide and propyteen oxide with thylenediamine) surfactants also available from BASE COD; and
- 6) compounds from (1), (2), (3) and (4) modified by "capping" or "end blocking" the terminal hydroxy 20 group or groups by reaction with small hydrophobic molecules such as propylece oxide, butylene oxide, benzyl choride, short chain fully acids, alrohols or ally! halides constaining from 1 to store 15 earthon alrons, 22 thionyl chloride, and so forth leading to all-block, 25 thionyl chloride, and so forth leading to all-block and block-heteric, betteric-block or all-block and so forth leading to all-block.

More suitably, the nonionics useful herein include, but are not limited to, block copolymers of ethylean oxide and propylene oxide sequentially condensed upon initiators hav-so ing diffunctional or tertrafunctional reactive hydrogens and alcohol alkoxylates. Especially preferred surfactants for compositions of the present invention are mixtures of slay in the property of the present invention are mixtures of slay in the propylene oxide sequentially condensed onto an ethylene-sate distance infiltation.

A blend of surfactants may be suitably employed in the present invention to arrive at the characteristics desirable for a particular application. For instance, some embodiments may include a surfactant for emulsification, a surfactant for soil removal, i.e. detersive surfactants, and so forth. Some embodiments may include the addition of a low foaming nonionic surfactants which have been found to be beneficial because they do not generate unwanted foam, do not interfere with antimicrobial activity, further solubilize otherwise insoluble or phase unstable fatty acids, and provide 45 improved surface wetting a solid penetration properties. Therefore, a blend of surfactants may be desirable. This part of the composition may therefore be referred to as the surfactant component to accurately reflect the fact that a single surfactant may be utilized in the compositions of the 50 present invention, or a blend including two or more surfactants may be utilized in the present invention. The surfactant component is generally useful from 0 wt-% to about 50 wt-% of the concentrate, suitably about 0.1 wt-% to about 50 wt-%, more suitably about 0.25 wt-% to about 45 wt-%, 55 even more suitably about 0.5 wt-% to about 40 wt-%, and most suitably about 1 wt-% to about 30 wt-% of the

As noted above, in some embodiments of the present invention, a couple or hydrotrope will suitably be added to the compositions, particularly when supplied in concentrated from to solubilize the falty acids in water. Those which have been found to be particularly effective for solubilizing the falty acids of the present invention include, but are not limited to, the anionic sulfornate surfactasts such as the alkali metal salts of C<sub>c,13</sub> sally sulfomates such as 64 10-ctane sulfonate, the alkali metal any 1 sulfonates, C<sub>c,30</sub> alkary) sulfonates such as the sulfact such as 54 10-ctane sulfonates, the sulfact install any sulfonates such as the sulfact such as 55 10-ctane sulfonates, the sulfact install any sulfonates such as the solution C<sub>c,30</sub> alkary applicates such as the solution C<sub>c,30</sub> alkary applicates.

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lene salfonates, sodium xylene salfonates, sodium cunene sulfonates, alkyl benzene sulfonates de sulfonyl had personate exten, and or forth Most sulfably, the sulfonyl benzene sulfably sulfonates alkyl benzene sulfably from shout 50 wt-% of the concentrate, sulfably from about 50 wt-% obout 50 wt-% on sulfably from about 15 wt-% to about 40 wt-%, and most sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% of the sulfably from about 5 wt-% to about 30 wt-% t

Commercially available bydrottops/couplets include, for crample, DOWAXW allystand diplernly oxide disaflorate surfactants; PETROL® allyl naphthalene sulforate surfactants; BIO-TEROS® PAS-8 cacen sulfonate surfactant sand so forth. The proportion of the surfactant component which is made up of a hydrotrope depends upon various factors including the specific hydrotrope cmployed, and the specific fatty acid employed, for instance. The hydrotrope is generally useful from 0% to about 50 wt % of the concentrate and suitably about 1 to about 40 wt % of the concentrate, and more suitably about 5 wt % to about 40 wt % of the concentrate.

The addition of an anionic hydrotrope has been found useful for maintaining product stability, and for decreasing the chance of phase separation over time.

The lists of ingredients given herein are intended as exemplary lists and are by no means exhaustive of the ingredients useful herein. Such lists are not intended to limit the scope of the present invention.

Other ingredients may be optionally added to the compositions of the present invention to impart additional propo crites to the composition in amounts which do not deterimentally affect the desired properties. Such properties may include form, function, aesthetics, and so forth. Such ingradients include, but are not limited to, solvents, other surfactants, couplers, defoamers, chelating agents, dyes, fregaraces, freeloogy modifiers, maunfacturing process aids, corrosion inhibitors, preserving agents, buffers, tracers, inert lillers and solidifying agents other antimicrobials, and so

The balance of the concentrates and/or diluted use solutions is typically water. A concentrate may or may not comprise any water. The concentrates may be diluted with any amount, but are typically diluted in the range of about 1:100 to about 1:1500 parts concentrate to water which are typical of normal use dilutions. For cleaning compositions, the compositions are typically more concentrated. For example, cleaning compositions may be diluted to ratios of about 1:100 to about 1:500, more suitably about 1:100 to about 1:300. For sanitizing compositions, the dilutions are typically greater than about 1:100 up to about 1:1500. A standard use dilution is about 1 ounce concentrate to about 6 gallons of water (2.957×10<sup>-2</sup> liters to about 3.785 liters or about 29.57 ml to about 3785.41 ml). This ratio is approximately 1:768 parts concentrate to water. The compositions may also be diluted with solvents other than water. However, water is the most commonly used solvent for dilution.

The compositions of the present invention may be prepared in various forms in both ready-to-use, and in concentrated versions. As noted above, the concentrated compositions require no dilution, but are typically formulated in one of several ways.

Commonly, the compositions are prepared as liquid comcentrates intended for further dilution just prior to use, or are prepared as ready-to-use compositions requiring no second distion. They may also be prepared as dispensable and dissolvable solid powders, tablets, blocks, or other solid matrix forming demicals well leavon to those of ordinary skill in the art. These examples are intended for illustrative purposes only. One of ordinary skill in the art understands that there are numerous modifications and other forms in which such compositions are available. Such modifications or changes in form can be made without departing from the scope of the present invention.

The compositions of the present invention have been 5 mond to be particularly suitable for use in cleaning and/or sanitizing operations because of their excellent stability at use dilutions, particularly in cooler water temperatures of 40–50° E (4.4–10° C). This property is particularly advantageous in food harvesting and flood and beverage processing operations to least din nod climate geographical regions

where water temperatures are often cooler. The present invention contemplates methods of using the composition for hard surface cleaning and/or sanitizing of in-place or clean-in-place (CIP)/SIP (steam-in-place) assemblies. The compositions may be introduced into a cleaning 15 and/or sanitizing system either manually, or using an automatic metering and/or dispensing system. The compositions may be either pre- or post-diluted with water before or after addition to the system. This is usually accomplished at ambient temperatures. The composition is then circulated 20 through the system, drained, and optionally, the system is rinsed one or more times with potable water. These CIP or SIP systems typically utilize low foaming compositions. However, high foaming compositions may be employed where foaming is not a concern and are contemplated as 25 being within the scope of the present invention as described above. For example, high foaming sanitizers may be employed for sanitizing external surfaces of equipment, ceilings, walls, floors, and so forth, while low foaming compositions may be employed for cleaning the internal equipment systems such as piping systems, i.e. dairies, for

The present invention also contemplates methods of using the compositions as one-step eleaner/santizers and disinfectants in which one composition can both clean and santitize a surface simultaneously. Typically the surface is <sup>35</sup> characterized as a hard surface. Such surfaces include equipment involved in both food and beverage processing such as in dairy operations including pipelines and bulk tanks and howaries.

Various modifications can therefore be made to the 40 present invention including modifications to the chemical formulation and to the physical form without departing from the scope of the present invention as described above.

The following non-limiting examples further illustrate the present invention.

# EXAMPLES

### Test Methods

1. Foaming Evaluation

Distilled water (300 mL) at a temperature of about 50–70° <sup>20</sup> E. was poured into a 500 mL granhard eylinder. Powdered product (10 g) or liquid product (10 mL) was poured into the graduated eylinder which was then stoppered tightly. The cylinder was then inverted and returned to an upright position 10 times. The graduated cylinder was then allowed 55 to sit and the water and form layers allowed to separate. The begint of the foat mayer in mL was determined at the highest and lowest points after the designated clapsed time. The average of the two readings was reported.

#### foam height=(mL of foam+liquid)-(mL of liquid)

2. Food Contact Surface Sanitizing Efficacy at 25° F.

Testing was conducted according to AÓAC Germicidal and Detergent Samitzing Action of Disinfectants 960.09, Official Methods of Analysis of the AOAC International, 65 16<sup>th</sup> Edition, 1995. Testing required EPA Pesticide Assessment Guidelines, Subdivision G 91-2(k)(Y2).

All of the examples tested were diluted at a ratio of 1 oz to 6 gallons concentrate to water (0.13%) using 500 ppm synthetic hard water (as CaCO). The hard water was prepared as follows:

#### Hard Water Preparation 500 ppm synthetic hard water (as CaCO<sub>3</sub>) PREPARATION

Solution A (mL)	Solution B (mL)	Total Volume (mL)	Final pH	Hardness Determination (ppm CaCO <sub>3</sub> )
- 5	4	1000	7.91	SDD nom

# Solution A:

31.74 g MgCl<sub>2</sub> (or equivalent of hydrates)+73.99 g CaCl<sub>2</sub> (or equivalent of hydrates) and diluted to 1 liter in boiled deionized water (heat sterilized) Solution B:

56.03 g NaHCO<sub>3</sub> diluted to 1 liter in boiled deionized water (filter sterilized)

Two test systems were used for this study per USEPA Pesticide Assessment Guidelines Subdivision G, Series 91, Subseries 91-A, 91-2, (k)(2).

# Staphylococcus aureus ATCC 6538 Escherichia coli ATCC 11229

% Reduction = Inoculum Numbers - Survivor Numbers × 100

Each of the following compositions was prepared by admixing listed chemicals in sequential order, blending thoroughly by agitation and allowing each ingredient to completely disperse or dissolve into liquid mixture before addition of the next ingredient. The resultant compositions were clear and homogeneously uniform upon admixture of all listed ingredients. The concentrates were conditioned at 40° F. until visual phase instability was observed or after 4 days with no visual change in stability. Use dilutions were prepared similarly using 1 oz of the concentrate per 6 gallons water (0.13%). The use diluted compositions were also conditioned at 40° F. for 4 days and observed for physical instability. The examples are illustrative of the stability results obtainable with compositions of the present invention. Variation was exhibited in the range, however, particularly in relation to the amount of time a composition remained at low temperatures. Stabilities varied and lesser stabilities were obtained depending on conditions, time, and composition.

## Examples 1-3

The following table 1 illustrates compositions of the present invention which utilize nitric acid as the strong acid and which contain no phosphoric acid.

TABLE

	1	2	3
Glacial acetic acid	15	15	15
Nitric acid. 42° Be	30	30	30
1-octane sulfonate, sodium, 40% active	25	25	25
Perlargonic acid (C <sub>0</sub> )	6	6	6
Decanoic acid (C <sub>10</sub> )	1	1	1
Urca	_	0.1	_
FD&C yellow #5, 0.10% (dyc)	-	0.44	_

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TABLE 1-continued

	1	2.	3	
DI water	23.00	22.46	23	- 5
Comment	CDS	dyc not stable	CDS	

CDS = concentrate and dilution stable, no visible precipitate/floc, very minor surface

# Examples 4-8

The following table 2 illustrates compositions of the present invention that utilize a blend of nitric acid and phosphoric acid and which contain no urea.

	TABLE	2				
	4	5	6	7	8	
Phosphoric acid, 75%	15	15	10	10	15	
Nitric acid, 42 deg Be	15	15	21	21	15	20
Glacial acetic acid	15	15	15	15	15	
1-octane sulfonate, sodium, 40% active	25	25	25	25	25	
Perlangonic acid (Ca)	6	6	6	2	7	
Decanoic acid (C <sub>10</sub> )	í	1	í	í	í	
FD&C Yellow #5, 0,10%	0.44	_	_	_		25
DI water	22,56	23	22	21	22	
Comments	dye not stable	CDS	CDS	CDS	CDS	

CDS = concentrate and dilution stable, no visible precipitate/floc, very minor surface

Examples 9-18

The following table 3 illustrates compositions of the present invention having a nitric acid/phosphoric acid blend and containing varying amounts of urea.

TABLE 3

9	10	11	12	13	14	15	16	17	18
10	10	10	10	10	10	10	10	10	10
21	21	21	21	21	21	21	21	21	21
15	15	15	15	15	15	15	20	20	20
25	25	25	25	25	30	25	25	30	30
7	7	7	7	7	7	6	7	7	6
1	1	1	1	1	1	1	1	1	1
0.10 20.9 CDS	0.50 20.5 CDS	1.00 20.0 CDS	5.00 16.0 CDS	0.25 20.75 CDS	0.5 15.50 CDS	0.5 21.50 CDS	0.5 15.50 CDS	0.5 10.50 CDS	0.5 11.50 CDS
	10 21 15 25 7 1 0.10 20.9	10 10 21 21 15 15 25 25 7 7 1 1 0.10 0.50 20.9 20.5	10 10 10 10 11 15 15 15 25 25 25 25 10 1.00 1.00 1.00 1.00 1.00 1.00 1.00	10 10 10 10 10 12 12 12 13 15 15 15 15 25 25 25 25 16 10 10 10 10 10 10 10 10 10 10 10 10 10	10	10 10 10 10 10 10 10 10 10 10 10 10 10 1	10	10	10

CDS = concentrate and dilution stable, no visible precipitate/floc, very misor surface

Examples 19-28

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The following tables 4-8 illustrate compositions of the present invention employing various surfactant blends.

TABLE 4

	19	20	21	22	23	24	25
Phosphoric acid, 75%	10	10	10	10	10	10	10
Nitric acid, 42 deg Be	21	21	21	21	21	21	21
Glacial acetic acid	15	15	15	15	15	15	15
1-octane sulfonate, sodium, 40% active	25	25	25	25	2.5	25	25
PLURAFAC® RA-40	1.00	_	_	_	_	_	_
TETRONIC ® 1307	_	2.00	_	_	_	_	_
TETRONIC © 1107	_	_	2.00	1.00	0.50	_	_
TETRONIC © 908	_	_	_	_	_	0.50	0.65
Perlargonic acid (Co)	7	7	7	7	7	7	7
Decanoic acid (C <sub>10</sub> )	1	1	1	1	1	1	1
Urea	0.5	0.5	0.5	0.5	0.5	0.5	0.5
DI water	19.50	18.50	18.50	19.50	20.00	20.00	19.85
Comments	ECDS						

ECDS = excellent concentrate and dilution stability, no visible precipitate/floc, no visible surface piling at 40° F.

PLURAFAC® RA-40 is an alcohol ethoxylate.

TETRONIC® 908, 1107 AND 1307 are all nomionic surfactants block copolymer adducts of ethylene oxide and propylene oxide to ethylenediamine.

25 Example 24 and Comparative Examples A-C Comparative Examples A-C are representative of com-mercially available sanitizing compositions which are stan-(Batch 3)

TABLE 5

dards in the industry.

Comparative A (wt-%)	Comparative B	Comparative C	
30% phosphoric soid	16% soft water	11% soft water	
21.99956% citric acid,	38% phosphoric acid,	35% phosphoric acid	
(50% active)	(75% active)	(75% active)	
9% citric acid, anhydrous	10% propicaic scid	8% lactic acid, food grade (88% active)	
30% 1-octane sulfonate, sodium (40% active)	3% perlargonic acid	34% sodium linear alkyl naphthalene sulfonate	
6% octanoic acid	3% decanoic acid	9% octanoic acid	
2% decanoic acid	30% 1-octane sulfonate, sodium (40% active)	3% decanoic acid	
q.s. isopropyl alcohol q.s. FD&C yellow #5	q.s. FD&C yellow #5		

A foaming evaluation was conducted according to Test Method #1. FIG. 1 is a bar graph showing the results of the foaming evaluation. As can be seen from the graph, example 50 24 exhibited a lower foam height than comparatives A-C which are standards in the industry.

Formula 24 was further tested for food contact surface sanitizing efficacy at 25° F. as described in Test Method #2 above. The following results were obtained.

TABLE 6 Efficacy Test Results

-	Staphylococcus aureus AFCC 6:	538
Test Substance (Batch Number)	Average CFU/mL of Test Survivors	Percent Reduction
Formula 24	3.0 × 10 <sup>1</sup>	>99.999
(Batch 1)	$1.8 \times 10^{2}$	>99.999
Formula 24	$7.5 \times 10^{1}$	>99.999
(Batch 2)	7.5 × 10 <sup>4</sup>	>99,999

TABLE 6-continued

	Efficacy Test Results Staphylococcus aureus ATCC 6	538
Test Substance (Batch Number)	Average CFU/mL of Test Survivors	Percent Reduction
Formula 24 (Batch 3)	5.5 × 10 <sup>1</sup> 2.8 × 10 <sup>2</sup>	>99.999 >99.999

# TABLE 7

	Efficacy Test Results  Escherichia coli ATCC 11229	_
Test Substance (Batch Number)	Average CFU/mL of Test Survivors	Percent Reduction
Formula 24	7.5 × 10 <sup>1</sup>	>99.999
(Batch 1)	<10	>99,999
Formula 24	<10	>99.999
(Batch 2)	<10	>99.999
Formula 24	<10	>99,999
(Batch 3)	<10	>99,999

As can be seen from tables 6 and 7, formula 24 exhibited a 99,999% reduction of S. aureus and E. coli. Example 24 therefore meets the efficacy requirements of a food contact surface sanitizer.

TABLE 8

	TEDENE O		
	26	27	28
Phosphoric acid, 75%	10	10	10
Nitric scid, 42 deg Be	21	21	21
Glacial acetic acid	15	15	15
1-octane sulfonate, sodium, 40% active	25	25	25
DOWFAX © C-6 acid,		_	_
DOWFAX ® C-6 acid, 78%	_	_	_
PETRO @ LBA liquid, 50%	-		-
TETRONIC @ 908	1.00	1.50	2.00
Perlargonic acid (C <sub>0</sub> )	7	7	7
5 Decanoic acid (C <sub>10</sub> )	1	1	1
Uren	0.5	0.5	0.5

TABLE 8-continued

	26	27	28
DI water	19.50	19.00	18.50
Commonts	ECDS	ECDS	ECDS

DOWFAX @ C-6 is a sodium hexyl diphenyloxide disulfonate PETRO LBA is a sodium alky naphthalene sulfonate

TETRONIC © 908 is a block copolymer adduct of ethylene oxide and amoviene oxide to ethylenediamine

TABLE 1

Cleaning Compositions		
	Example 29	Example 30
Phosphoric acid, 75%	20.0	20.0
Nitrie Acid, 42 Be	21.0	21.0
Glacial Acetic Acid	15.0	15.0
1-octane sulfonate	10.0	20.0
TETRONIC © 908	0.5	1.50
Perlargonic acid	1.0	3.4
Decanoic acid	0.15	0.5
Ures	0.5	0.5
DI water	26.85	18.10

The above compositions are illustrative of compositions of the present invention which are useful as cleaning compositions, i.e. one-step cleaning compositions. Example 29 is intended for 1% dilution (1:100 concentrate to water) and example 29 is intended for 0.3% (1:333 concentrate to 30 water) dilution.

What is claimed is:

- A sanitizing and/or cleaning composition comprising: a) about 3 wt-% to about 12 wt-% of the composition of at least one aliphatic short chain antimicrobially effective C5 to C14 fatty acid or mixture thereof;
- b) about 5 wt-% to about 50 wt-% of the composition of at least one weak C1 to C4 carboxylic acid; and
- c) about 5 wt-% to about 50 wt-% of the composition of a strong acid which is nitric acid or a mixture of nitric and phosphoric acids
- 2. The composition of claim 1 wherein said at least one weak carboxylic acid is acetic acid, hydroxyacetic acid, propionie acid, hydroxypropionie acid, alpha-ketopropionie acid, citric acid, butyric acid, valeric acid, succinic acid, tartaric acid, malic acid, fumaric acid, adipic acid, formic acid, or mixture thereof
- 3. The composition of claim 1 wherein said at least one aliphatic short chain antimicrobially effective fatty acid is a
- C<sub>6</sub> to C<sub>10</sub> fatty acid.

  4. The composition of claim 1 wherein said at least one 50 weak carboxylic acid is acctic acid.
- 5. The composition of claim 1 wherein said aliphatic short chain fatty acid is decanoic, nonanoic or a mixture thereof.
- 6. The composition of claim 1 wherein said aliphatic short chain fatty acid is nonanoic acid. 7. The composition of claim 1 wherein said aliphatic short
- chain fatty acid is present at a concentration of about 5 wt-% to about 10 wt-% of the composition. 8. The composition of claim 5 wherein said mixture comprises about 0.25 to about 5 wt-% of the composition
- decanoic acid and about 2 to about 10 wt-% of the composition nonanoic acid. 9. The composition of claim 5 wherein said mixture comprises about 0.5 to about 4 wt-% of the composition
- decanoic acid and about 3 to about 9 wt-% of the composition nonanoic acid. 10. The composition of claim 5 wherein said mixture comprises about 1 to about 3 wt-% of the composition

16 decanoic acid and about 4 to about 8 wt-% of the compo-

- sition nonanoic acid 11. The composition of claim 5 wherein said mixture comprises about 1 wt-% of the composition decanoic and about 6 to about 7 wt-% of the composition nonanoic acid.
- 12. The composition of claim 1 wherein said at least one weak carboxylic acid is present at a concentration of about 10 wt-% to about 40 wt-% of the composition.
- 13. The composition of claim 1 wherein said at least one weak carboxylic acid is acetic acid.
- 14. The composition of claim 1 wherein said phosphoric acid is present at a concentration of about 5 wt-% to about 40 wt-% of the composition.
- 15. The composition of claim 1 wherein said phosphoric acid is present at a concentration of about 10 wt-% to about 35 wt-% of the composition.
- 16. The composition of claim 1 wherein said nitric acid is present at a concentration of about 5 wt-% to about 50 wt-%
- of the composition. 17. The composition of claim 1 wherein said nitric acid is present at a concentration of about 15 wt-% to about 40 20 wt-% of the composition.
  - 18. The composition of claim 1 further comprising an effective amount of urea to reduce nitrogen peroxide to
- 19. The composition of claim 18 wherein said effective amount of urea is about 0.05 wt-% to about 5 wt-% of the 25 composition.
  - 20. The composition of claim 18 wherein said effective amount of urea is about 0.5 wt-% of the composition. 21. The composition of claim 1 further comprising at least
  - one surfactant 22. The composition of claim 21 wherein said at least one
  - surfactant is nonionic. 23. The composition of claim 22 wherein said at least one surfactant is a tetrafunctional block copolymer derived from the addition of ethylene oxide and propylene oxide to
  - ethylenediamine. 24. The composition of claim 21 wherein said surfactant is present at a concentration of 0.1 wt-% to about 50 wt-%
  - of the composition. 25. The composition of claim 1 further comprising at least one organic hydrotrope
- 26. The composition of claim 25 wherein said organic hydrotrope is an anionic sulfonate or corresponding disulfonate 27. The composition of claim 26 wherein said organic
- hydrotrope is an alkyl sulfonate, an aryl sulfonate, a alkaryl sulfonate or a corresponding disulfonate, an alkylated diphenyl oxide disulfonate, or an anionic mono or disubstituted alkyl ethoxylated phosphate ester, or a mixture thereof
- 28. The composition of claim 27 wherein said organic hydrotrope is 1-octane sulfonate.
- 29. The composition of claim 25 wherein said organic hydrotrope is present at a concentration of about 0.5 wt-% to about 50 wt-% of the composition.
- 30. The composition of claim 1 further comprising water. 31. The composition of claim 30 wherein said concentrate is diluted with water at a ratio of 1:100 to about 1:1500 parts
- concentrate to water 32. The composition of claim 30 wherein said concentrate is diluted at a ration of about 1:768 parts concentrate to
- 33. A method of one-step cleaning and sanitizing a surface comprising the step of contacting said surface with a composition as in claim 1.
- 34. The method of claim 33 further comprising the step of diluting said composition with water at a ration of about 1:100 to about 1:1500 of the composition to water.
- 35. A clean-in-place method of cleaning a beverage or food processing unit including conduits, surfaces and containers, comprising the steps of:

- a) providing the composition of claim 1; and
   b) contacting conduits, surfaces and containers in said beverage processing unit,
- c) removing said composition from said unit for the purpose of reinitiating processing.

36. The method of claim 35 further comprising the step of diluting said composition with water at a ratio of about 1:100

to about 1:1500 of the composition to water.

37. A dilutable acid sanitizing and/or cleaning concentrate composition comprising

about 0.25 wt-% to about 10 wt-% of the concentrate of at least one C<sub>6</sub> to C<sub>10</sub> fatty acid or mixture thereof;

about 5 wt-% to about 50 wt-% of the concentrate of a C<sub>1</sub> to C<sub>4</sub> weak carboxylic acid;

about 0 wt-% to about 40 wt-% of the concentrate phosphoric acid;

about 5 wt-% to about 50 wt-% of the concentrate nitric acid; and

about 0.05 wt-% to about 5 wt-% of the concentrate urea; 20 with the proviso that the concentration of nitric acid and phosphoric acid does not exceed about 50 wt-% of the concentrate.

38. The composition of claim 37 wherein said phosphoric acid is present at a concentration of about 5 wt-% to about 25 40 wt-% of the concentrate.

39. The composition of claim 37 further comprising at least one surfactant at a concentration of about 0.1 wt-% to

about 50 wt-% of the concentrate.

40. The composition of claim 37 further comprising 30

water.
41. A cleaning composition comprising:

a) about 0.5 wt-% to about 10 wt-% of at least one short chain fatty acid:

b) 0 wt-% to about 40 wt-% phosphoric acid;

c) about 5 wt-% to about 40 wt-% phosphoric acid;

d) about 0.5 wt-% to about 50 wt-% of at least one surfactant; and

e) about 5 wt-% to about 50 wt-% of the concentrate of a  $_{40}$ 

C<sub>1</sub> to C<sub>4</sub> weak carboxylic acid; with the proviso that the concentration of nitric acid and phosphoric acid does not exceed about 50 wt-% of the concentrate.

- 42. The composition of claim 41 wherein said composition comprises about 5 wt-% to about 40 wt-% of said at least one surfactant.
- 43. The composition of claim 42 wherein said at least one surfactant is aniomic.
- 44. The composition of claim 43 wherein said at least one surfactant is an alkyl sulfonate, an aryl sulfonate, a  $c_{a,30}$  alkaryl sulfonate or a corresponding disulfonate, an alkylated diphenyl oxide disulfonate, or an anionic mono or disubstituted alkyl ethoxylated phosphate ester, or a mixture thereof
- 45. The composition of claim 44 wherein said at least one surfactant is a 1-octane sulfonate.
- 46. The composition of claim 41 wherein said at least one surfactant is nonionic.
- surfactant is nonionic.

  47. The composition of claim 46 wherein said at least one surfactant is a tetrafunctional block copolymer derived from
- the addition of ethylene oxide and propylene oxide to ethylenediamine.

  48. The composition of claim 46 wherein said at least one
- surfactant is present at a concentration of about 0.25 wt-% to about 10 wt-%.

  49. The composition of claim 41 wherein said at least one
- fatty acid is present at a concentration of about 1 wt-% to about 5 wt-%.

  50. The composition of claim 41 wherein said at least one fatty acid comprises about 0.1 wt-% to about 5 wt-%
- decanoic acid and about 0.5 wt-% to about 10 wt-% nonanoic acid.

  51. The composition of claim 41 wherein said at least one fatty acid comprises about 0.1 wt-% to about 1 wt-% decanoic acid and about 1 wt-% to about 5 wt-% nonanoic
- acid.
  52. A sanitizing and/or cleaning composition comprising:

a) 1 part of at least one aliphatic short chain antimicrobially effective C<sub>2</sub> to C<sub>1.4</sub> fatty acid or mixture thereof; b) about 1.7 to about 4.2 parts of at least one weak C<sub>1</sub> to C, carboxylic acid; and

 about 1.7 to about 4.2 parts of a strong acid which is nitric acid or a mixture of nitric and phosphoric acids.

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